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- (54) New catalyst components for the polymerization of ethylene and of mixtures thereof with olefins and catalysts obtained therefrom.
- (5) Components of catalysts for the polymerization of ethylene and of mixtures thereof with olefins, in the form of emulsions or dispersions in an inert liquid medium or in inert gas phase of a liquid phase comprising a compound or composition containing a compound of the metals of the Groups IV to VI of the Periodic System, immiscible in aliphatic hydrocarbons or which components are obtained from emulsions or dispersions in an inert tiquid medium or in gas phase of a precursor of the catalyst component, which in the liquid state is immiscible with the normal aliphatic hydrocarbons.

COMPLETE DOCUMENT

Object of this invention are new catalyst components for the polymerization of ethylene and of the mixtures thereof with olefins and the catalysts obtained therefrom.

More particularly, the invention refers to new liquid catalytic components used in emulsion or dispersion in an inert liquid medium and the use of the catalysts deriving therefrom in the polymerization of ethylene and of mixtures thereof with alpha-olefins and/or with polyenes containing at least two double bonds.

It is known that the coordination catalysts commonly used in the industrial practice are heterogeneous systems obtained by reaction of a transition metal compound (generally a Ti halide) with an organometal compound of the metals of Groups I to III of the Periodic System.

The transition metal compound used for the preparation of the catalyst is generally a solid insoluble in the hydrocarbon polymerization medium or is a liquid soluble in said medium.

Homogeneous coordination catalysts (soluble in the polymerization medium at least before the polymerization is started) are also known.

These systems, however, have not been adopted in the industrial practice because of the fact that the activity thereof, that at the beginning is also very high, decreases rapidly and this does not allow to obtain high polymer yields.

Within the field of heterogeneous catalysts, supported

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ready since some time, which are endowed of so high activity as to allow to avoid the expensive depuration treatments of the polymer from the catalytic residues. These catalysts are generally obtained from a catalyst component comprising a titanium compound supported on a magnesium halide in active form.

In the modern industrial processes using "high yield" supported catalysts the requirement is felt of having available catalysts with controlled morphology and particle size, capable to yield a polymer in form of particles reproducing the morphology and the particle size of the catalyst and furthermore endowed with high flowability and bulk density.

A catalyst having these characteristics shows advantages not only during polymerization step and makes easier the subsequent operations of transfer and/or treatment of the polymer but can also allow to avoid the granulation step of polymer. This operation, as it is known, requires large amounts of energy.

The methods employed until now for the preparation of catalysts having a controlled morphology and/or particle size are based on operations expensive per se, which comprise the preformation of a precursor of the catalytic component in form of particles having a controlled morphology and the subsequent transformation of these particles to obtain the true catalyst component or the precipitation of

the catalyst component under conditions in general very critical, followed by the steps of separation, washing and dry ing of the solid.

According to other methods, precursors of the catalyst component in the melted state are emulsified in an inert im miscible liquid and the emulsion is then subjected to quanching to obtain the solidification of the dispersed liquid phase which is subsequently treated for the transformation the reof to catalyst component.

Examples of these methods are described in the applicant's U.S. patent 3,953,414 and Belgian patent 878,347.

Coordination catalysts are not known hitherto wherein the component comprising the transition metal compound in the form of a liquid immiscible in the conventional polymerization solvents, is employed in emulsion or dispersion in said liquid medium.

It has been now unexpectedly found that it is possible to obtain polymers of ethylene and copolymers thereof with olefins in the form of particles having controlled morphology and/or particle size, by carrying out the polymerization with catalysts obtained from emulsions or dispersions in an inert liquid medium or in an inert gas phase of a liquid phase immiscible in the liquid medium, comprising a compound or compositions containing a transition metal compound of the Groups IV to VI of the Periodic System, that are immiscible, in the liquid state, with the normal aliphatic hydrocarbons.

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The catalyst components of the invention consist in particular of emulsions or dispersions in an inert liquid medium or in an inert gas phase of a liquid phase comprising compounds or compositions containing compound of Ti, V, Zr and Cr, in particular compounds of Ti, that are immiscible in the normal aliphatic hydrocarbons (hexane, heptane and the like). Within the scope of the invention fall also the liquid or solid catalytic components obtained by post-treatment of the catalytic emulsions or dispersions of the invention.

Also the solid or liquid catalyst components form part of the invention, that are obtained by subjecting to transformation post-reactions in order to obtain the true catalytic component emulsions or is persions in an inert liquid medium or in an inert gas phase of precursors of catalyst components, the precursors being immiscible, in the liquid state, in the normal aliphatic hydrocarbons.

According to another aspect of the invention the transition metal compound immiscible in aliphatic hydrocarbons or the liquid precursor are made to be adsorbed on solid porous supports having a controlled morphology and/or particle size: the support is then used in suspension in an inert liquid medium immiscible with the adsorbed liquid phase or dispersed in gas phase.

Examples of solid inert supports are silica, alumina and polymers in form of porous particles having controlled morphology and or size.

As already indicated, any transition metal compound of the Groups IV to VI that is immiscible in the liquid state with aliphatic hydrocarbons is suitable for the preparation of the emulsions or dispersions forming the catalytic components of the invention.

Similarly, any precursor of the catalytic components which is immiscible, in the liquid state, with the aliphatic hydrocarbons may be used to prepare the emulsions or dispersions useful for the preparation of the catalytic components of the invention, such as the reaction product of an anhydrous magnesium halide, in particular MgCl₂ of a Mg dialkyl or a Grignard reagent, with an anhydrous aluminum halide, in particular AlCl₃, in an aromatic hydrocarbon in particular toluene, in the presence of a halogenated hydrocarbon, preferably 1,2-dichloroethane. To the oily liquid thus obtained (non immiscible in the same aromatici hydrocarbon), a titanium compound, in particular TiCl₄ is added. An oily liquid is obtained that is immiscible in the normal aliphatic hydrocarbons.

More particularly, the Mg and Al halides and the aromatic hydrocarbon are caused to react at the reflux temperature of the hydrocarbon in the Mg/Al/toluene molar ratio of 1:2:12.

To this suspension the halogenated hydrocarbon is added in the ratio of about 2 mols per mol of Mg halide and it is heated until an oily liquid is formed. Then the Ti compound is added in a Ti halide/Mg halide comprised in generale between 0.1:1 and 1:1.

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In the above preparation the halogenated hydrocarbon can be replaced by anhydrous gaseous hydrochloric acid or by a normally liquid halogenated transition metal compound. For example TiCl₄ is particularly suitable to this purpose. In the latter case no further addition of the transition metal compound is needed to prepare the catalyst component.

According to another embodiment of the invention the Mg dihalide can be added last and solubilized in the mixture obtained by first mixing AlCl₃ and toluene and by thereafter treating the mixture with the halogenated hydrocarbon or gaseous HCl or TiCl₄.

The oily liquid that is obtained before the addition of the Ti compound is in itself a precursor of the catalytic components that can be emulsified and the emulsion can be treated for the transformation thereof to the catalyst component.

In the above indicated preparation the Al trihalide can be substituted by an alkyl Al-dihalide, the toluene by benzene, xylene and by similar aromatic hydrocarbons; 1,2-dichloroethane can be substituted by C_2H_5Cl , C_3H_7Cl , $n-C_4H_9Cl$, $s-C_4H_9Cl$, $t-C_4H_9Cl$, C_6H_5Cl , $CHCl_3$, $C_6H_5CH_2Cl$, CH_2Cl_2 and by alike alkyl, aryl or arylalkyl halides.

Another method of preparation consists in dissolving an anhydrous Mg halide in a Ti tetraalcoholate, in particular Ti tetrabutylate, and in flowing an anhydrous gaseous hydrohalogenic acid through the solution until an oily phase is separated. According to a modification of the above describ

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ed method, the hydrohalogenic acid can be replaced by an acyl chloride, in particular acetyl chloride. The butyl acetate that is formed is partly removed until formation of an oily liquid.

Other compounds suitable for the preparation of the catalytic components of the invention can be selected from the compounds having the formula:

that in the liquid state are immiscible with aliphatic hydrocarbons. In the formula X is a halogen atom; Y is a OR radical in which R is an alkyl, cycloalkyl or aryl group containing from 1 to 18 carbon atoms; p is a number from 1 to 4; q is a number from zero to 3; p+q = 4; n is a number from 3 to 6; ED is an electron-donor compound selected in particular from the esters of carboxylic aliphatic or aromatic acids.

Examples of these compounds are: MgCl₂.TiCl₄.4 Ac (AC = = CH₃COOC₂H₅), MgCl₂.TiCl₄.4 EB (EB = ethylbenzoate), MgCl₂.TiCl₂(OC₂H₅)₂.4C₂H₅OH; MgCl₂.TiCl₄.5POCl₃.

These compounds are prepared according to known methods, by dissolving the Mg halide in the ED compound, by adding to the solution the stoichiometric amount of the Ti compound and then by causing the reaction to take place at reflux. The excess of the ED compound is at the end removed by evaporation.

As already indicated, the compounds or the compositions

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containing the transition metal compounds, which form the emulsions or dispersions of the invention, can be subjected to modification or transformation reactions before that the emulsions or dispersions are employed as catalyst component. Said reactions comprise the reactions with electron-donor compounds, in particular esters, Al-alkyl compounds and silicon compounds.

The emulsion is then made to react, according to known methods, with reagents capable to transform the adduct in active catalytic component.

Known transformation reactions to catalyst component are those wherein the adduct is made to react with ${\rm TiCl}_4$ or with an Al-alkyl compound or with halogenated silicon compounds such as ${\rm SiCl}_4$ and halosilanes. Examples of these reactions are described in the Belgian patents 857,574 and 878,347.

As already indicated, the oily liquid obtained from the reaction of MgCl₂, AlCl₃, toluene and dichloroethane can be utilized as precursor and made to react in emulsion or dispersion with substances capable to transform it to the catalyst component. For example it can be reacted with TiCl₄ and alkyl hydropolysiloxanes, according to known methods.

The emulsifying of the transition metal liquid compounds as well as that of the precursors is carried out according to known techniques.

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The liquid compounds can be dispersed in the same hydrocarbon medium used in polymerization, optionally in the presence of surfactants of known type selected from those that are inert or poorly reactive toward the catalyst components. In the processes in gas phase, the transition metal compound can be dispersed directly in the gas phase containing the co-catalyst.

Preferably, however, the transition metal compound is dispersed previously into an oil of parafinic, naphthenic, aromatic or siliconic type; the resulting dispersion or emulsion is then fed to the polymerization reactor containing the co-catalyst.

Examples of these oils are silicon oil Baysilon M 100 (Bayer), vaselin 0 55 (Rol), Cortis M 100 oil (Total), Circosol 2XH oil (Sunoco) and Dutrex R55 oil (Shell).

The addition during the emulsifying of an electron-donor compound and/or of the co-catalyst in amounts in general comprised between 5 and 50% by weight on the amount of the transition metal compound has beneficial effects on the morphologyc characteristics of polymer.

In some cases, in particular when the emulsion are unstable in the time or would create feeding problems, it has been found convenient to prepolymerize small amounts of ethylene or other olefin. The prepolymerization is carried out in general until a few grams of polymer per gram of catalyst component are formed.

A solid stable prepolymer is obtained having spherical

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and regular size particles which subsequently during the pole lymerization maintain or reproduce the shape while growing, thus generating polymer particles that are also regular and spherical.

The co-catalysts to be utilized together with the catalytic components of the invention are the organometal compounds of Al, preferably the not halogenated compounds, as for example $\mathrm{Al}(\mathsf{C_2H_5})_3$, $\mathrm{Al}(\mathrm{iC_4H_9})_3$, $\mathrm{Al}(\mathsf{n-C_4H_9})_3$, $\mathrm{Al}(\mathsf{C_4H_9})_2^H$, $\mathrm{Al}(\mathsf{C_6H_{13}})_3$. The polymerization of ethylene or of mixtures thereof with alpha-olefins with these catalysts is carried out according to any of the known techniques, in the presence or the absence of hydrocarbon media, in liquid or gas phase, at temperatures preferably comprised between 0° and 200 °C and by operating in general in the presence of molecular weight regulators of polymer, as hydrogen.

Al already indicated, the catalysts are used in the polymerization of both ethylene and of mixtures thereof with alpha-olefins, C_3 - C_{12} , in particular butene, to obtain crystalline copolymers of ethylene containing minor amounts of polymerized alpha olefins as well as to obtain elastomeric ethylene-alpha-olefin copolymers, in particular ethylene-propylene copolymers containing optionally insaturations derived from polyenes having at least two double bonds.

In using the emulsified or dispersed catalyst component of the invention, precautions have to be taken to introduce the catalyst component into the polymerization reactor in the form of the emulsion or dispersion as prepared. Means

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to avoid the demixing of the dispersed or emulsified liquid phase are well known in the art. For instance, the emulsified or dispersed catalyst components should be introduced into the polymerization reactor under the polymerization conditions as quickly as possible, immediately after its preparation.

As mentioned at the beginning, the advantages obtainable with the use of these catalysts are multiple. These catalysts, besides being endowed of activities in general very high (such as to not require any purification treatment of the obtained polymer from the catalytic residues), show the advantage of furnishing polymer powders endowed of excellent morphological characteristics, high bulk density and very restricted particle size distribution. Consequently, the polymers possess a so high flowability and bulk density that they can be utilized directly in the conventional operations of molding and extrusion, without having recourse previously to a costly grapulation operation.

Another advantage of remarkable interest is the extreme semplicity of preparation of these catalysts, which reduces considerably the cost thereof, also with respect to other known, highly active catalysts.

The melt index E and respectively N were determined according to the ASTM methods D-1238 condition E and respectively N, measured at 190 °C and expressed in g/10 minutes.

The tamped bulk density was determined according to a method described in DIN 53194.

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The particle size was determined according to ASTM D 1921-63.

The flowability and the poured bulk density were determined according to ASTM D-1895-69.

The following examples are given to illustrate the invention without limiting it.

Examples 1-8

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Preparation of catalytic component A.

The catalyst component A used in all tests of examples 1-8 was prepared in the conditions and with the modalities as follows.

9.5 g (0.1 mol) of anhydrous powdered magnesium chloride (water content lower than 1% by weight), 39.9 g (0.3 mol) of anhydrous aluminum trichloride, and 128 cc of anhydrous toluene were introduced in said order into a 500 cc flask fitted with mechanical stirrer, dropping funnel, reflux condenser and thermometer, that was previously flushed with nitrogen.

The resulting toluene suspension was heated to 110 °C and maintained under stirring at this temperature for 2 hours. The mixture was then cooled to 45 °C and 15.7 cc of 1,2-dichloroethane were added thereto with the dropping funnel.

After heating at 100 °C for 1 hour, a red-brownish liquid was obtained, to which, after cooling again to 45 °C, 0.1 mol of $TiCl_A$ was fed.

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After an additional 15 minutes stirring at 45 °C, the solution obtained results to have a volume of 178 cc and the following composition:

Ti = 2.35% by weight

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Al = 4.5% by weight

C1 = 22.75% by weight

Mg = 1.2 % by weight

The viscosity at 20 °C was 3.5 centistokes. The density at 20 °C was 1.13 g/cc.

Dispersion of catalytic component A. The apparatus for the dispersion or emulsifying of the ca-

talytic components that was used in all examples consists of a small turbostirrer (Ultra Turrax TP 18/10 manufactured by Janke & Kunkel, Ikawerk, Staufen) turning at a speed of 12,000 r.p.m. that was inserted in a 150 cc Keller type flask fitted with a 50 cc graduated dropping funnel and a thermometer. The control of the temperature was carried out by immersion of the flask in a methanol /dry ice cooling system.

Prior to use, the apparatus was flushed with nitrogen. The dispersing oil, the optional additives coadjuvant for the dispersion were introduced in this order into the flask, and the catalytic component was then introduced in the time of 5 minutes under the higheest stirring while simultaneously cooling and maintaining the temperature at 20 °C. The stirring was then continued for another 10 minutes.

At the end of the dispersion operation and after reducing to 50% the stirring speed, the emulsion obtained was injected in the polymerization autoclave in the amounts reported in Table 1.

Polymerization of ethylene

The polymerization test of all examples was carried out in a 2.5 liter stainless steel autoclave fitted with an elicoidal blade stirrer operating at the speed of 600 r.p.m. and with an automatic thermostatic system.

At the moment of introducing the catalytic dispersion the autoclave duly flushed with nitrogen contains already 1 liter of anhydrous hexane and 7.5 mMols of Al-triisobutyl that are maintained under stirring at 60 °C. After introduction of the catalytic component in the hexane, the temperature was raised to 75 °C, 3 atm. of $\rm H_2$ (partial pressure) were introduced and lastly the autoclave was pressurized with ethylene up to the total pressure of 14 atmospheres.

The autoclave was then stirred for 2 hours while maintaining constant the pressure by continuous feeding of ethylene. The feed of ethylene was discontinued, the reactor was cooled to room temperature, the polymer was discharged and separated from hexane by filtration and then dried in nitrogen flow at 70 °C for 8 hours.

The amounts of the reactants used in the preparation of the dispersions of the catalytic component in the different types of oil and the polymerization results are reported in Table 1.

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In Example A of Table 1 are reported the reuslts of the polymerization of ethylene with the catalytic component A as such (not subjected to emulsifying).

Examples 9-15

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Preparation of the catalytic component B

The catalytic component B utilized in examples 9-15 was prepared under the conditions and with the operating modalities as follows.

- 15 0.2 mols of anhydrous powdered magnesium chloride and 0.4 mols of titanium tetrabutylate were introduced into a 1000 cc flask fitted with mechanical stirrer, reflux condenser, thermometer and dropping funnel. The flask was heated at 140 °C for 3 hours.
- The resulting solution, cooled to 60 °C, was diluted with 640 cc of anhydrous n-heptane. Then gaseous anhydrous hydrochloric acid was introduced by means of a glass tube immersed directly in the liquid phase at 20 °C and under stirring, in such amount as to maintain the temperature of the system at about 30 °C.

During this treatment, there takes place first a turbidness of the solution and then the separation of an oily reddish phase immiscible in heptane.

After 3 hours the feed of hydrochloric acid was interrupted and the oily suspension was heated while stirring at 60 °C for 30 minutes. Said liquid suspension was transferred at room temperature to a dropping funnel and then the oily

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phase was separated. 206 cc of oily liquid were obtained which had the following composition by weight:

Ti = 7.4 %

Mg = 1.75 %

Cl = 22.15 %

 $C_4 H_0 OH = 45.7 \%$

The viscosity at 20 °C is of 180 cst and the density at 20 °C is of 1.056 g/cc. For the dispersion or emulsifying of the catalytic component B in the different types of oil and for the experimenting with it, including the polymerization tests therewith, the same operating modalities already described for the catalytic component A were adopted. The results were reported in Table 2.

In Example B of Table 2 the results of polymerization of ethylene using the catalytic component B as such (not dispersed) were reported.

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Table 1

E	ΞX.	AMPLE Nº	Α	1	S	3	4	5	6	7	8
COMPONENT	NT.	Туре	-	BAYSILON	BAYSILON M 100	BAYSILON N 100	BAYSILON M 100	VASEL IN	CORTIS M 100	CIRCOSOL 2XH	DUTREX
	S AGENT			oil	oil	oll	011	ol 1	oi 1	oil	R 55 oll
CATALYTIC	SPERSING	Viscusity at 20 °C(cst)	-	140	140 .	140	140	55	500	91,5	175 (100 °C)
2	PISF	Quantity (cc)	-	20	15	20	20	20	20	20(1)	20(1)
N OF THE	ADDITIVE	Туре	-	ı	-	n.butyl ether	4mM A](Iso butyl)+n, butyl ather	-	-	-	
BS I		Quantity (cc)	•	•	-	3	3	-	-	-	-
BISPERSION		atalytic liquid A	•	20	30	20	20	20	10	10(1)	10(1)
	٢	ispersed catalyst sed (cc)	0.3	0,5	0, 15	0,3	0.3	0.3	0,1	0, 15	0,3
	۳	eight polymer obtained	239	135	198	205	183	121	185	270	104
	Ϋ́	-3 eld(g.pol./g.Tl).10	30	20	75	55	49	30	209	203	39
	и	eli Index E (g/10')	0,21	0.4	0.45	O _• 5	0,21	0,25	0,35	0,23	0,14
.T.S	Me	elt Index E	10.7	8.1	9.2	8	7.6	11.6	8,3	8.9	7.2
RESULTS	de	amped bulk ensity (g/cc)	0,25	0.39	0,27	0,41	0,43	0.27	0.35	0.23	0, 18
		Soher'ical polymer	NO	YES	YES	YES	YES	YES	YES	МО	YES
ZATION		Particle form polymer	но	NO	но	NO	NO	NO	YES	YES	но
POLYMERIZATION	אטע געופט איי פארא איים		YES	ю	NO	NO	но	YES	NO	но	YES

(1) Dispersion carried out at 50 °C.

		EXAMPLE N°	В	9	10	11	12	13	14	15
	AGENT	Туре	-	BAYSILON M100 oll	BAYSILON H 100 oil	VASELIN OB 55 oll	CORTIS 100 M oll	CIRCOSOL 2XH oll	DUTREX R 55 oll ·	n-HEPTA- NE
GONDAID	PS ING	Viscosily at 20 °C (ast)	-	140	140	55	500	91,5	17.5 (100°C)	0.57
Й	810	Quantity (cc)	-	20	10	20	20 .	20	20	50
DISPERSION OF THE CATALYTIC COMPONENT	IVE	Type	-	-	Al(n-oc tyl)3	-	-	-	-	FENOROL 10
RS I ON OF	AD01TIVE	Quantity (cc)	-	-	3.7 (m.mol§)	-	-	-	-	0.03 (g)
SPE	С	atalytic liquid B (cc)	-	20	20	10	10	10	10	10
	۵	ispersion used (cc)	0,01	0,01	0.02	0.05	0.03	0.05	0.05	0.02
	Wo	ight polymer obtained (g)	154	91	178	123	254	268	91	103
	YI	eld (g.pal./g Ti) x 10 ⁻³	197	233	171	95	299	206	70	3'96 `
ŀ	Мо	alt Index E (g/10')	0,25	0,3	0.25	0.15	0, 18	0.23	0.2	0,12
	I —	olt Index N	8	8.1	7.8	8.0	10	7.4	8	7.5
ESUL TS	T	amped bulk density (g/cc)	0,35	0.33	0,35	0,22	0.24	0.28	0,23	0.21
I S	>	Spherical polymer	МО	МО	YES	YES	YES .	YES	YES	YES
POLIMERIZATION RESULTS	MORFOLOSY	Particle form polymer	, NO	YES	МО	YES	YES	но	YES	YES
Por	POLYMEP M	I iii equiti porymet	YES	YES	но	ОИ	NO	, YES.	но .	NO

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Examples 16 and 17

Preparation of the catalytic component C.

The catalytic component C utilized in examples 16 and 17 was obtained in the following manner.

- 10 0.2 mols of anhydrous powdered MgCl₂ and 0.16 mols of titanium tetrabutylate were introduced in this order into a 250 cc flask fitted with mechanical stirrer, reflux condenser and thermometer.
- The flask was heated while stirring the suspension at 140 °C for 3 hours. A milky liquid was obtained, to which, after cooling to room temperature, 1.2 mols of acetyl chloride were added in 30 minutes. The obtained suspension while still under stirring was heated at boiling (73 °C) for 2 hours.
- Then the unreacted acetyl chloride was removed by distillation. The temperature in the flask is let raise up to 125 °C. A this point the residue has the appearance of an oily red-brownish mass.
- It amounts to 90 cc and has a density of 1.25 g/cc at 20 °C.

 The results of the polymerization tests were reported in

 Table 3. Test C-was carried out using the catalytic component C not dispersed.

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Table 3

		EXAMPLE N°	С	16	17
THE CA-	AGENT	Туре	-	BAYSILON M 100 OIL	CORTIS M 100 OIL
ON OF THE	SING	Viscosity at 20 °C (cst)	- .	140	500
RSION IC CO	10-	Quantity (cc)	••	20	20
DISPERSION TALYTIC COM	Ca	talytic liquid C (cc)	-	10	10
====		alytic dispersion used (cc	0.2	0.3	0,2
	Weig	ht polymer obtained (g)	287	. 182	230
S		d (g.pol/g Ti).	16,000	20,000	38,000
RESULTS		Index E (g/10')	0.07	0.07	0.07
RES	Mel	Index N t Index E	10.2	9.7	9.3
TION	Tam	ped bulk dens <u>i</u> (g/cc)	0.23	0.27	0.28
ZI ZA	MOR-	Spherical polymer	NO	YES	YES
POLYMERI ZATI ON	MER MC	Particle form polymer	NO	YES	YES
POI	POLYMER	Irregular polymer in powde	r YES	NO	NO

Example 18

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The following substances were used for the preparation of the dispersion of the catalytic component:

Baysilon M 100 oil = 30 cc

n-butyl ether = 4.5 cc

Catalytic component A = 30 cc

The resulting dispersion was introduced into a 2.5 l reactor containing already 1000 cc of anhydrous hexane and 0.1 mol of aluminum triisobutyl at the temperature of 40 °C. Ethylene was then introduced till a pressure of 0.6 atm. The polymerization starts immediately and causes a temperature increase to 45 °C.

While maintaining constant said conditions, ethylene was continuously fed for 30 minutes, i.e. until about 3 g of ethylene are polymerized per gram of catalytic component A. The pre-polymerized catalytic component was then discharged and, after removal of the solvent by filtration, it was subjected to two washings with anhydrous hexane at room temperature.

The catalytic component in hexane suspension was then used in subsequent tests of ethylene polymerization, in order to evaluate the possible variations in performance with respect to the time (ageing).

In table 4 the results of said tests (ageing of the prepoly merized catalysts: zero days; 5 days; and 13 days) were reported.

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Furthermore, some morphological characteristics of the polymers obtained (flowability and particle size distribution) were reported.

In all cases the powders obtained consisted wholly of spherical particles.

In the case of the polymer obtained with aged catalyst, the analysis of the catalytic residues furnished the following results:

Ti = 2 ppm

Cl = $\langle 25 \text{ ppm} \rangle$ Mg = 4.6 ppm

Total ahses = 0.095%

The polymerization test was carried out in the conditions of examples 1-8 with the difference that the duration is of 4 hours in place of 2 hours.

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Table 4

_	eing of the prepolyme- zed catalyst	zero days	5 days	13 days
Yi	eld (g polymer/g Ti).	313,000	340,000	315,000
Ме	lt Index E (g/10')	0.23	0.14	0.2
	lt Index N lt Index E	8	8.3	8.3
Та	mped bulk density (g/cc)	0.4	0.41	0.4
Fl	lowability (sec.)	17.5	17	18
y WEIGHT)	> 1000 M (and (2000 M))	25.1	30.4	30.2
ION (% BY	> 500. Ju	53.3	52.1	54
DISTRIBUTION	> 177 µ	19.7	16.7	15
SIZE	> 105 µ	1,6	0.5	0.5
PARTICLE	< 105 µ	0.3	0.3	0.3

Example 19

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300 g of polyethylene obtained according to example 18 (ageing time of 5 days) were introduced into a 3 liter stain less steel autoclave fitted with an anchor-shaped stirrer. After heating at 80 °C for 1 hour under a flow nitrogen and under stirring at intervals, the polymer was subjected to treatment with 0.5 g of aluminum triisobutyl diluted in 500 g of propylene. The mixture was stirred at 70 °C for 2 hours and then all propylene was removed slowly.

15 0.634 g of prepolymerized catalyst (obtained as described in example 18) and then, at the temperature of 60 °C, 1 g of aluminum triisobutyl dissolved in 20 cc pentane were charged into the reactor.

The reactor was heated at 75 °C and while maintained under stirring was charged with 1.8 atm. of hydrogen and then with ethylene up to a total pressure of 9 atm. The pressure was maintained constant during the polymerization by continuous addition of ethylene.

After 4 hours, the reactor was degassed and the dry polymer was collected which weighed 578 g. The yield was 231,000 g polymer/g titanium used.

The morphology of the obtained polymer was wholly spherical.

The tamped bulk --- density was 0.38 g/cc. The flowability was 18 seconds.

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The particle size distribution was as follows:

 $\emptyset > 1000 \,\mu (\text{and } < 2000 \,\mu) = 33.5 \,\%$ by weight $\emptyset > 500 \,\mu$ = 59.4 % by weight $\emptyset > 177 \,\mu$ = 6.6 % by weight $\emptyset > 105 \,\mu$ = 0.4 % by weight $\emptyset < 105 \,\mu$ = 0.1 % by weight

Example 20

Two polymerization tests were carried out, one in hexane and the other in isobutane, and both were carried out in the presence of butene-1 in different amount.

In the two tests it was utilized a prepolymerized catalyst from the same preparation used for the polymerizations reported in Table 4, eccept that it had 90 days of ageing.

The autoclave used was entirely similar to that of the preceding polymerization, except that it had a 1.5 l capacity in place of 2.5 l.

Polymerization in hexane

Anhydrous hexane containing Al-triisobutyl and the weight catalyst were charged into the autoclave duly flushed with nitrogen.

The reactor was heated at 70 °C and then flowed with hydrogen and pressurized with the same up to 3 atmospheres. Simultaneously 18 g of butene-1 and ethylene up to a total pressure of 14 atmospheres were introduced. The pressure was maintained constant by introducing continuously ethyle-

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ne and butene, the latter one in an amount of 0.6 g per each 30 g of ethylene fed.

After 2 hours polymerization, the reactor was degassed and cooled by stripping with steam. The polymer was recovered from the slurry and then dried at 70 °C in a nitrogen flow for 12 hours.

Polymerization in isobutane.

20 cc hexane containing the catalyst and Al-triisobutyl were charged into the flushed autoclave wherein a flow of isobutane is maintained. Then also isobutane in liquid state was introduced. The reactor was heated at 70 °C (isobutane pressure = 10 atm.). 4 atmospheres of hydrogen were introduced therein and simultaneously 180 g of butene-1 and ethylene up to a total pressure of 29.5 atmospheres. The polymerization was continued in these conditions for 2 hours, by adding 2.95 g of butene-1 per each 30 g of ethylene fed. The reactor was degassed and cooled to room temperature. The polymer was recovered and dried at 70 °C in a nitrogen flow for 12 hours.

The data concerning both tests and the results were reported in Table 5.

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.<u>Table 5</u>

	Polymerization in	hexane	isobutane
Hex	ane (cc)	700	20
lso	butane (g)	·	220
A1-	triisobutyl (g)	1	1
Pre	polymerized catalyst (g)	0.3	0.625
But	ene-1 (g) { total feed	18	180
		13.2	194.75
1	ained polymer (g)	65	175
1 .	eld (g/g Ti)	114,000	147,000
Mel	t lndex E (g/10') .	0.17	1.6
	N/E	10.6	7.6
	F/E	31.2	25.6
Bor	nded butene (% by weight)	4 2	4.5
Dei	sity (g/cc)	0.9394	0.9262
Bu:	lk density (Tamped) (g/cc)	0.31	0.33
Flo	owability of the polymer (sec.)	23	24.4
	Morphology	wholly spherical	wholly spherical
	>2000 µ (% by weight)	0.9	1.4
E	>1000 µ (% by weight) .	26.1	23 •
SIZE	> 420 M (% by weight)	52.9	57.2
	> 250 µ (% by weight	12.8	12.3
E	> 149 µ (% by weight)	5.3	4.4
ric	>105 µ (% by weight)	1.2	1.0
PARTICLE	< 105 M (% by weight	0.8	0.7

Excluding silicon oil, the carbon of the chemical structures of the dispersing oils used in examples 5 and 11, respectively 6 and 12; 7 and 13; and respectively 8 and 14 is distributed percentwise as follows:

Oil	Aromatic	Naphthenic	Paraffinic
	carbon	carbon	carbon
Vaselin OB 55 Cortis 100 M Circosol 2XH Dutrex R 55	-	-	100
	7	28	65
	20	39	41
	49	35.5	17.5

Example 21

magnesium chbride were introduced in this order into a 200 cc Keller type flask fitted with a turbostirrer of the type used in examples 1-8 and of a dropping funnel. 4.35 cc of anhydrous ethanol were added slowly to the mixture. The su spension was heated slowly to 80 °C and then to 125 °C after inserting on the Keller a reflux condenser in order to stop the evolution of ethanol vapours.

The emulsifying of the adduct MgCl₂.alcohol that is formed, was obtained by vigorous stirring. 300 cc of ISOPAR G and 8.25 cc of TiCl₄ were introduced into a glass vessel with

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jacket for the circulation of a thermostatic liquid, and fitted with mechanical stirrer, reflux condenser and thermometer.

The temperature of the solution was increased to 120 °C and at said temperature the whole volume of the emulsion obtain ed as indicated was introduced therein slowly (about 5 minutes).

The whole system was maintained at 120 °C for 1 more hour. After removing the liquid by siphoning, the solid was washed with hot ISOPAR G. A sample containing 0.004 g of solid catalytic component was taken from the suspension of the last washing and it was used in a polymerization test of ethy lene that was carried out in the conditions described in examples 1-8.

50 g of polymer (yield of 12,500 g/g catalytic component) were obtained in the form of particles having a controlled particle size (most of the particles have the size comprised between 500 and 1000 micron).

Example 22

Example 21 was repeated except that the emulsion of the adduct of MgCl₂ and ethanol was added to a solution of 8.3 cc of TiCl₄ in 300 cc of ISOPAR G that is maintained at 120 °C. At about a half way of addition of the emulsion, another 8.3 cc of TiCl₄ were added, and the addition of the emulsion was then completed.

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-.31 .--

In a polymerization test carried out in the conditions of example 21 and using 0.0055 g of catalytic component, 120 g of polymer were obtained in the form of spheroidal particles having the size mainly comprised between 500 and 1000 micron.

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CLAIMS

1) Components of catalysts for the polymerization of ethy lene and of mixtures thereof with olefins, in form of emulsions or dispersions in an inert liquid medium or in inert gas phase of a liquid phase comprising a compound or a composition containing a transition metal compound of the Group IV-VI of the Periodic System, immiscible with aliphatic hydrocarbons, or that are obtained by subjecting to transformation reactions forming the catalyst component, emulsions or dispersions in an inert liquid medium or in inert gas phase of a precursor of the catalyst component, that is immiscible with aliphatic hydrocarbons.

2) Catalyst components according to claim 1, in which the transition metal compound is selected from the compounds of Ti, V, Zr and Cr.

- 3) Catalyst components according to the preceding claims, in which the dispersed liquid phase comprises compounds of Ti and Mg.
- 4) Catalyst components according to claim 3, in which the dispersed liquid phase containing the Ti and Mg compound is obtained by reacting an anhydrous halide of Ti and Mg and an aluminum halide or alkyldihalide in an aroma-

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- tic liquid hydrocarbon, in the presence of a halogenated hydrocarbon or by reacting an anhydrous halide of Mg and a Ti alcoholate, in the presence of an anhydrous hydrohalogenic acid.
 - 5) Catalyst components according to claim 3, in which the Ti and Mg compound has the formula:

 ${
m MgX}_2.{
m TiX}_p{
m Y}_q.{
m nED}$ wherein X is a halogen; Y is a radical OR (R = Alkyl, cycloalkyl or aryl group containing from 1 to 18 carbon atoms); p is a number from 1 to 4; q is a number from 0 to 3; p+q = 4; n is a number from 3 to 6; ED is an electron-donor compound.

- 6) Catalyst components according to the preceding claims, in which the compound or composition containing the transition metal compound is subjected to reactions and/or to modifying treatments before the use of the emulsion or dispersion as catalyst component.
- 7) Catalyst components according claim 6, in which the emulsion or dispersion is reacted with compounds selected from the Al-alkyl compounds, preferably in the presence of an olefin or from the silicon compounds and the electron-donor compounds.

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- 8) Catalyst components according to claim 1, in which the precursor of the catalytic component is a compound or a composition containing a Mg compound.
- 9) Catalyst components according to claim 8, in which the Mg compound is selected from the adducts of a Mg halide with an electron-donor compound, preferably an aliphatic, cycloaliphatic or alkylaryl alcohol.
- 10) Catalyst components according to claims 8 and 9, in which the emulsion or dispersion of the precursor of the catalytic component is reacted with a compound selected from the Ti halides, the Al-alkyl compounds and the halogenated Si compounds.
 - 11) A catalyst component according to claim 1, in which the aliphatic hydrocarbon-immiscible liquid compound of the transition metal is adsorbed on solid porous supports having a controlled morphology and/or particle size.
- 12) Catalysts for the polymerization of ethylene and of the mixtures thereof with alpha-olefins which comprise the reaction product of an organometal compound of Al and a catalyst component according to preceding claims 1 to 11.

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A	EP-A-0 018 737 (*Claims 1-12*	MITSUI)		1				
A	FR-A-2 428 056 (*Claims 1-6; exam)	1				
A	US-A-4 218 339 I CUFFIANI) *Claims 1-21; 6 52-68*	•		1				
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